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84-109393/an
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** SS 1: Results 1

Search statement 2

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1/1 DWPI - (C) Thomson Derwent

AN - 1984-109393 [18]

XA - C1984-046221

TI - Manganese, cobalt and nickel solutions extraction via solvents - consists of acid treatment, neutralisation, re:extraction, and regeneration of organic solution

DC - E31 J01 M25

PA - (SCHO/) SCHORTMANN P C

NP - 2

NC - 2

PN - ES8401143 A 19840216 DW1984-18 *

AP: 1982ES-0517148 19821105

- PT--77614 A 19840921 DW1984-44

PR - 1982ES-0517148 19821105

IC - C22B-023/04 C22B-047/00

AB - ES8401143 A

Process comprises treatment with mixts. of hydroxy-oximes and carboxylic acids, the acidity being neutralised with alkali, and the cobalt and nickel extract going for washing and reextraction using controlled pH acid. The organic extract containing nickel goes for reextraction giving an aqueous solution of nickel salt.

- A fraction of the organic solution, prior to recycling, undergoes regeneration in contact with an element in the metal state and then with an acid

with an acid.

MC - CPI: E35-S E35-V E35-W J01-K01 M25-G11 M25-G17 M25-G19

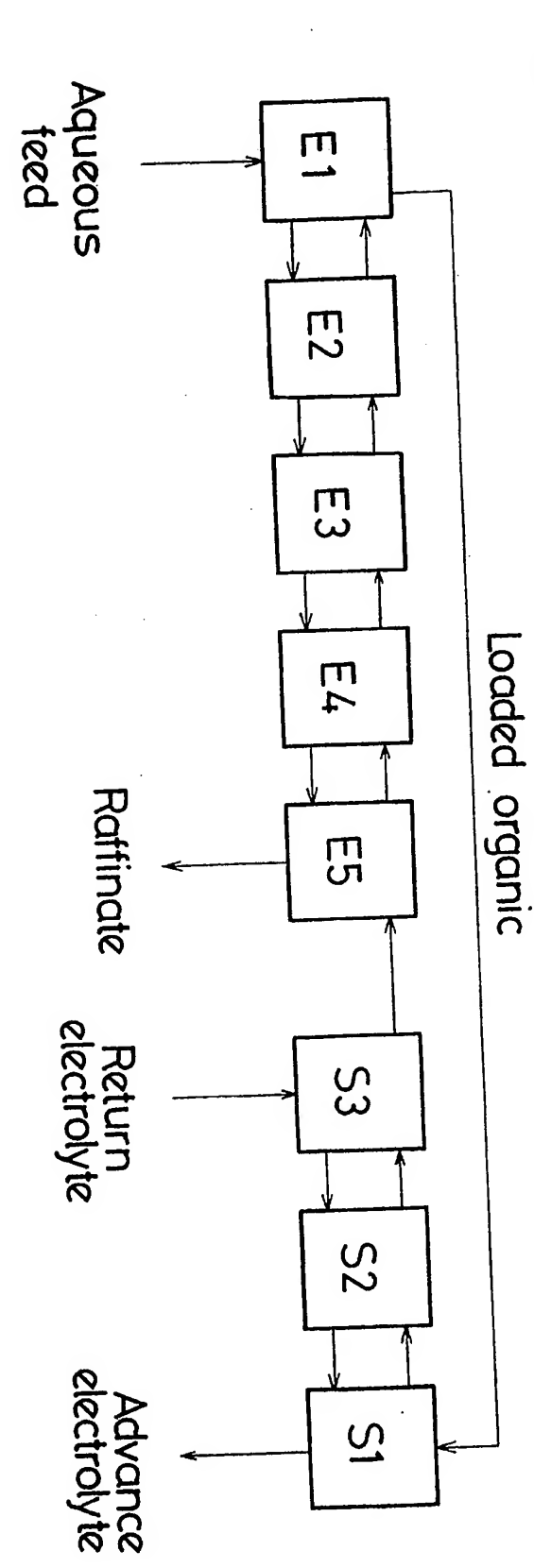
UP - 1984-18

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 - (71) Applicant Council for Mineral Technology (South Africa) 200 Hans Strydom Avenue Randburg Transvaal Province Republic of South Africa
 - (72) Inventor John Stephen Preston
 - (74) Agent and/or Address for Service D Young and Co 10 Staple Inn London WC1V 7RD

- (54) Selective solvent extraction using organophosphorus and carboxylic acids
- (57) A process for the extraction of metal values, in particular divalent metal ions such as copper, nickel, zinc, cobalt and manganese from aqueous solutions thereof which may be well in the acid range for example of pH 2 to 4. Extraction is effected using known organophosphorus or carboxylic acid extractants but in combination with nonchelating oximes, preferably of aldehydes wherein the alpha-carbon atom is primary or secondary. A synergistic effect between the extractant and oxime enables extraction to take place at a lower pH than heretofore thereby saving in acid and neutralizing agents in many instances.

TWO SHEETS - SHEET ONE



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TWO SHEETS - SHEET TWO

SPECIFICATION

Selective solvent extraction using organophosphorus and carboxylic acids

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5	BACKGROUND TO THE INVENTION THIS INVENTION relates to selective solvent extraction using organophosphorus and carboxylic acids as the metal extractants in the presence of additives which greatly enhance the utility of	5
10	extractants under certain conditions. It is often desirable to remove, and possibly recover, metals which are dissolved in a leaching step in hydrometallurgical processes. Metals which are often to be removed and recovered include cobalt, nickel, copper, zinc and manganese which are all amenable to removal by use of a solvent extractant.	10
15	However, as a result of the fact that leach liquors containing these metals usually result from a leach using acid, the pH of the leach liquor is generally rather low. Conventional solvent extractants such as organophosphorus and carboxylic acids only operate effectively at appreciably high pH and thus, either extremely poor performance must be tolerated, or the leach liquor must be at least partly neutralized thereby consuming both acid and alkali in order to	15
20	achieve this. Once at least partially neutralized, the organophosphorus i.e. organophosphoric, organophosphonic and organophosphinic, and carboxylic acids operate effectively as solvent extractants. Usually, however, such a procedure is not favoured as it is uneconomical and it would be most desirable to provide a process for the extraction of such metals at a substantially lower pH than is at present possible.	20
25	It is known, and has been practiced, to employ di-(2-ethylhexyl) phosphoric acid at very mildly acidic pH. The phosphonic acid equivalent has also been employed with somewhat greater success and a phosphinic acid reagent has recently become available. The above-mentioned phosphoric acid has also been used together with chelating additives in the form of the commonly known hydroxyoxime sold under the Registered trade mark LIX63 by Henkel Corporation for separating copper and cobalt from nickel in sulphate streams. This process did	25
30	not proceed, as far as Applicant is aware, beyond the pilot plant scale.	30
35	BRIEF SUMMARY OF THE INVENTION It has now surprisingly been found that an unforeseen extremely marked synergistic effect is obtained when there is used, together with an organophosphorus acid and any carboxylic acids having required extraction characteristics, a non-chelating oxime. This synergistic effect enables the metals to be extracted at substantially lower pH levels than was possible heretofore. Accordingly this invention provides a process for the solvent extraction of metal values from	35
40	solutions containing such metals comprising the contacting of a pregnant solution with a solvent extractant being an organophosphorus acid or a carboxylic acid capable of extracting desired metal values, the extraction being characterised by the fact that it takes place in the presence of a non-chelating oxime selected to enhance extraction of the desired metal values. Further features of the invention provide for the oxime to be selected to enable extraction to	40
45	take place at a lower pH than was practical heretofore; for the extractant to be organophosphoric acid of the type (RO) ₂ PO-OH where each R is, independently of the other, a substituted or unsubstituted alkyl, aryl, or alkaryl group; for the extracted metals to be cobalt, nickel, manganese, copper and zinc; for the extraction, in the case of organophosphoric acids to be carried out in the pH range of from 0 to 5, and in the case of carboxylic acids in the pH range	45
50	of from 1 to 5, and for the contact time to be of the order of a few minutes. The fact that a synergistic effect is created by employing non-chelating oximes together with the organophosphorus or carboxylic acids is evidenced by the fact that, in the case of cobalt and nickel, the order of extraction is entirely reversed so that the selectivity for cobalt over nickel is reversed. Also the extracted species are different: in the case of di-(2-ethylhexyl) phosphoric acid	50
55	with 2-ethylhexanal oxime the cobalt is extracted as the bright pink octahedral complex as opposed to the dark blue tetrahedral complex whilst the nickel is extracted as the turquoise as opposed to the apple-green octahedral complex in which the oxygen-donor ligands have been replaced by nitrogen-donor ligands.	55
	Not only does the present invention provide for extraction of metal values from solutions at a substantially lower pH than was heretofore possible, but the extraction takes place extremely quickly (of the order of a few minutes, as opposed to some hours in the case of a carboxylic acid and chelating oxime such as LIX 63 mixture) and, in addition, stripping of the extracted values from the solvent extractant has proved to be extremely simple. One reason for this is the avoidance of the formation of intractable cobalt (III) chelate complexes which are not amenable to conventional stripping techniques. Thus, stripping can be effected simply by using dilute	60
65	mineral acid solutions for example 20 to 200 g/l sulphuric acid.	65

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Also, some mixtures, for example 2-bromo octanoic acid mixed with 2-ethylhexanal oxime show useful separations of nickel which is extracted at pH 2 to 3 from cobalt which is extracted at pH 3 to 4 as opposed to the non-selective extraction of both metals together at a pH of 4 to 5 in the case of the carboxylic acid alone.

Preferably, the oximes employed in the implementation of the invention are those of aldehydes having the alpha-carbon as a primary or secondary one. Other oximes show similar synergistic effects.

It will be understood that the organophosphorus acids which can be used to extract cobalt, nickel, manganese and copper in the pH range from 0 to 2 enables these metals to be recovered directly from acid leach liquors without the necessity of partially or totally neutralizing the residual acid concentration of the leach liquors. This represents a substantial saving of both acid and neutralizing agents.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a series of graphs showing percentage extraction vs pH for four different extractant 15 systems and,

Figure 2 is a schematic flow sheet of a continuous counter current extraction process.

DETAILED DESCRIPTION OF VARIOUS EMBODIMENTS OF THE INVENTION

In order to investigate and confirm the operation of the invention initial tests were carried out under fixed conditions in order to determine the pH at which 50 per cent extraction would take place (pH 0,5) in the case of nickel and cobalt. These results are given in the Table below with each result having been obtained by investigating the extraction of cobalt and nickel by 0,5M organophosphorus acid in xylene at 20°C. In each case where an additive was added the extent of addition was also 0,50M. In each case the aqueous solutions contained 0,10M cobalt (II) and/or 0,10M nickel as the nitrate in 1,00M ammonium nitrate solution. The organophos-

phoric, organophosphonic and organophosphinic acids were investigated, the latter two being commercially available reagents RD 577 marketed by Shell Chemicals and Cyanex CNX marketed by the American Cyanamid Company in the case of the phosphonic and phosphinic acid reagents respectively.

35				рН _{0,5}		Δ p $H_{o.}$	5††		
30	Extractant	Additive†	Со	Ni	Nì — Co	Со	Ni	3	
40	Di-(2-ethyl- hexyl) Phosph- oric Acid		•						
	(D2EHPA)	None TBP	3,68 3,82	4,11 4,07	0,43 0,25	0,00 - 0,14	0,00 0,04	2	
		isodecanol nonylphen-	3,65	3,86	0,21	0,03	0,25		
45		ol TOPO	3,70 3,90	3,91 4,10	0,21 0,20	- 0,02 - 0,22	0,20 0,01	4	
	RD577	LIX 63 EHO None	0,12 1,99 3,90	ND* 1,58 5,12	ND . – 0,41 1,22	3,56 1,69 0,00	ND 2,53		
50		TBP isodecanol	3,99 4,12	4,98 4,77	0,99 0,65	- 0,09 - 0,22	0,00 0,14 0,35	Ę	
	CNX	EHO None**	3,25 4,22	3,06 5,20	- 0,19 0,98	0,65 0,00	2,06 0,00		
55		EHO**	4,22	4,58	0,36	0,00	0,62	-	

†TBP = Tri-n-butyl phosphate; TOPO = tri-n-octylphosphine oxide; EHO = 2-ethyl-hexanal oxime

 $\dagger \dagger \Delta p H_{0.5} = p H_{0.5}^{\text{without additive}} - p H_{0.5}^{\text{with additive}}$

*not determined due to extremely slow extraction rate of nickel

60 "solutions include 25 volume % isodecanol

It will be noted that only small influences on the pH_{0,5} were obtained in all cases apart from those in which 2-ethylhexanal oxime and LIX63 were employed. Whilst a vast improvement was obtained in the case of LIX63 with the organophosphoric acid in respect of cobalt, the extraction of nickel was too slow to be of any practical use and thus, at the best, this additive could be

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employed for separating cobalt from nickel. However, it must be borne in mind that extraction of cobalt even is slow and also stripping thereof from the reagent is extremely difficult.

In all other cases nickel pH recovery is substantially improved and an examination of the pH figures given in the cobalt and nickel columns shows that the oxime additive has by far the most desirable effect at lowering the pH value at which extraction takes place. This is, of course, with the exception of LIX63 the disadvantages of which are discussed above.

The pH extraction curves were plotted for various systems and exemplary curves are illustrated in Fig. 1. In Fig. 1, the curve numbered 1 is that of the synergistic mixture of 0,5M Di-(2-ethylhexyl) phosphoric acid and 0,5M 2-ethylhexanal oxime.

Curve numbered 2 is that of 0,5M RD573 (a commerically available chelating di-oxime made by Shell Chemicals) and only approaches the performance of the synergistic mixture of the invention. This chelating di-oxime has the disadvantage indicated above in that intractible cobalt complexes will arise unless a high selectivity over cobalt is exhibited.

Curve numbered 3 is that of 0,5M LIX65N (a variant of LIX63 and also made by Henkel 15 Corporation).

Curve numbered 4 is that of 0,5M Di-(2-ethylhexyl) phosphoric acid itself.

The curves were obtained using an aqueous phase of 0,1M Ni (NO₃)₂ in 1,0M NH₄NO₃ and the organic solvent was xylene. The temperature was 20°C, aqueous to organic ratio 1:1 and the contact time 10 minutes except in the use of LIX65N where the contact time was 60 minutes.

The superiority of the synergistic mixture of the invention is quite obvious from the curves which show that extraction takes place over the range of about pH 1 to pH 2, for the mixture.

Different Oximes

Tests were conducted to show that different oximes of the non-chelating type exhibited the same synergistic effect. In these tests the pH_{0,5} was again determined from the same 0,1M nitrate solutions employed in the previous tests. The aqueous to organic ratio was 1:1 and the organic phase was a 0,5M solution of oxime plus 0,5M Di-(2-ethylhexyl) phosphoric acid. The temperature was 20°C.

The oximes employed were all of the non-chelating type having the general formula R.CNOH.R' in which R and R' had the meanings set out in the following Table 2:—

TABLE 2
Extraction of nickel and cobalt by mixtures of Di-(2-ethylhexyl) phosphoric acid and non-chelating oximes, R.CNOH.R'.

				рН _{0,5}		Δρ	H _{0,5}	
40	R	R'	Ni	Со	Co-Ni	Ni	Со	40
40	C_7H_{15} $C_4H_9CH(C_2H_5)$	H H	1,34	1,72	0,38 0,41	2,77 2,53	1,96 1,69	40
45	(01.3/30	H H H	1,73 3,41 3,46	2,13 3,25 3,50	0,40 0,16 0,04	2,38 0,70 0,65	1,55 0,43 0,18	45
	C ₆ H ₁₃ C ₅ H ₁₁ C ₆ H ₅	CH₃ C₂H₅ CH₃	3,80 3,94 4,03	3,49 3,59 3,71	- 0,31 - 0,35 - 0,32	0,31 0,17 0,08	0,19 0,09 0,03	
50	(ČH₃)₂CH	(CH ₃)₂CH	4,15	3,77	- 0,38	- 0,04	- 0,03	50

Useful synergistic effects (i.e., large $\Delta pH_{0,5}$ values) are shown for the oximes of aldehydes where the alphacarbon is primary or secondary. Other oximes (namely those of aldehydes where the alpha-carbon is tertiary, aromatic aldehydes, and aliphatic or aromatic ketones) show very much smaller synergistic effects. Note that a positive value in the column headed $pH_{0,5}$ (Co-Ni) indicates that the mixture is selective for nickel over cobalt, whilst a negative value indicates selectivity for cobalt over nickel.

Different Organophosphoric Acids

Tests were conducted under the same conditions as for the preceding set of tests but with different organophosphoric acids and 2-ethylhexanal oxime as the non-chelating oxime. The different acids used were those having the following formulae:—

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The results are set forth in the following Table 3:-

Acid No. (Ig)

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TABLE 3
Extraction of nickel and cobalt by mixtures of organophosphoric acids and 2-ethylhexanal oxime.

	Organophos- pH _{0,5} for acid alone		eid	pH _{0,5} for acid plus oxime			ΔpH _{0,5}		5	
10		Ni	Со	Ni-Co	Ni	Со	Co-Ni	Ni	Co	
10	la lb	4,11 3,72	3,68 3,42	0,43 0,30	1,58 1,30	1,99 1,70	0,41 0,40	2,53 2,42	1,69 1,72	1
	lc ld	2,92 2,71	2,85 2,50	0,07 0,21	0,84 0,66	1,23 0,98	0,39 0,32	2,08 2,05	1,62 1,52	
15	le If	2,30 2,63	2,30 2,53	0,00 0,10	0,81 0,42	1,18 0,79	0,37 0,37	1,49 2,21	1,12 1,74	1

 $\Delta pH_{0.5} = pH_{0.5}$ for acid alone - $pH_{0.5}$ for acid plus oxime.

These results show that by varying the molecular structure of the organophosphoric acid, in particular by introducing aromatic functions, extraction of cobalt and nickel can be carried out under remarkably acidic conditions and that extractant mixtures can be formulated for the relevant circumstances.

It is to be noted that the selectivity of the organophosphoric acids for cobalt over nickel (column 4) becomes one of nickel over cobalt for the mixed extractants (column 7).

Different Carboxylic Acids

Tests were conducted on carboxylic acids with 0,5M carboxylic acid (1,0M for Versatic 10) plus, where applicable, 0,5M non-chelating oxime in xylene (one result labelled‡ used heptane as the diluent). The aqueous phase contained 0,1M Ni (NO₃)₂ or 0,1M Co(NO₃)₂ in 1,0N NH₄NO₃. Contact was made at aqueous to organic ratio of 1:1.

The results obtained are reflected below:—

TABLE 4
35 Extraction of nickel and cobalt by mixtures of carboxylic acids and non-chelating oximes.

	Carboxylic	Non-chelating		рН _{0,5}		Δр	H _{0,5}	
40	acid	oxime†	Ni	Co	Co-Ni	Ni	Co	40
	Versatic 10	None	5,98	6,16	0,18	0,00	0,00	
	Versatic 10	CAO	4,22	5,01	0.79	1,76	1,15	
	Versatic 10	EHO	4,66	5,28	0,62	1,32	0,88	
45	2-bromolauric	None	4,68	4,70	0,02	0,00	0,00	45
		EHO	2,56	3,47	0.91	2,12	1,23	70
	2-bromocapry- lic	CAO	2,34	3,30	0,96	ND	ND	
		EHO	2,42	3,61	1,19	ND	ND	
50		EHO‡	1,96	3,20	1,24	ND	ND	50
		EBO	2,70	3,70	1,00	ND	ND	30

†CAO, caprylaldehyde oxime; EHO, 2-ethylhexanal oxime; and EBO, 2-ethylbutanal oxime.

Versatic 10 is a highly branched carboxylic acid produced by Shell Chemicals.

Although mixtures of carboxylic acids and non-chelating oximes do not permit extraction of cobalt and nickel under as acidic a condition as mixtures of organophosphoric acids and non-chelating oximes, the nickel-over-cobalt selectivity of the former mixtures (column 5) are significantly higher than the latter (column 7 in Table 2) and the required pH is certainly lower.

Effects on Different Cations

Tests were carried out on numerous different cations, generally metal ions, in order to ascertain the effect of the non-chelating oxime. The tests were conducted, as before, by determining the pH_{0,5} in each case. All cations were present as nitrates in 1M HNO₃ and the pH 65

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was adjusted using concentrated ammonia solution. The exceptions to the above were the ferrous Fe^{2+} ion which was in the sulphate form in HNO_3 initially and the Cr^{2+} and VO^{2+} cations which were in the sulphate form in $0.33M\ H_2SO_4$ the pH of which was adjusted using NaOH.

The extractant was 0,5M Di-(2-ethylhexyl) phosphoric acid (D2EHPA) and 0,5 M2EPHA in admixture with 0,5M 2-ethylhexanal oxime in xylene in each case and at 20°C. Contact took place at an aqueous to organic ratio of 1:1.

The results of the tests are reflected in Table 5 below:-

10	Metal	pH _{0,5} (D2EHPA)	pH _{0.5} (D2EHPA-EHO)	ΔpH _{0,5‡}	10
	Ni ²⁺	4,12	1,60	2,52	
	Cu ²⁺	2,90	1,05	1,85	
15	Co ² +	3,70	2,00	1,70	4.5
	Fe ^{2 +}	3,56	2,10	1,46	15
	Mn ²⁺	2,90	2,12	0,78	
	Cr²+	2,15	1,47	0,68	
	VO ² +	1,25	0,96	0,29	
20		7	-,	0,20	20
	Fe³+	-0.32	-0,40	0,08	20
	Cr ³⁺	3,12	2,98	0,14	
	La³+	1,96	2,06	- 0,10	
25	Mg²+	3,81	3,69	0,12	2.5
	Ca ² +	2,85	2,97	-0,12	25
	Zn²+	1,42	1,58	-0.16	
	Cd ²⁺	2,98	1,48	1,50	
	Ag+	3,25	< 0,00	> 3,25	
30	Pb ² +	2,50	2,33	0,17	
	NH +	3,57*	3,37*	0,20	30

 $$\pm \Delta pH_{0.5} = pH_{0.5}(D2EHPA) - pH_{0.5}(D2EHPA-EHO)$$

*pH at which the NH + concentration in the organic phase

35 reaches 0,05M.

Appreciable synergistic effects are shown by all the diavelent transition metal ions studied (note that VO²⁺, however, has only one 'vacant' coordination site once complexed with D2EHPA, in contrast to two for the other divalent metal ions). There is no synergistic effect for trivalent ions.

40 Amongst the non-transition metals, only silver and cadmium show significant synergistic effects. The implications for extraction of the divalent transition metals without co-extraction of gangue metals (magnesium, calcium, etc) or neutralization products (e.g. ammonium ion) are clear.

Continuous Counter-Current Extraction

Based on information collected a counter current type of extraction process was tested using a Bell-Engineering mixer-settler apparatus. The test extraction was that of nickel as it showed excellent sensitivity to the non-chelating oxime synergistic effect. The test work which will not be described in detail herein resulted in good extractions of Ni taking place over five extraction stages E1 through E5 as shown in Fig. 2. Three stripping stages are employed.

Typical operating conditions were found to be as indicated below in the case of use with and without the oxime. The system tested had organic phases consisting of 15% D2EHPA and 15% D2EHPA together with 5% EHO in Solvesso 150 as solvent. (Solvesso is an aromatic type of solvent sold under Trade Mark "Solvesso 150" by Esso Chemicals.

The organic to aqueous ration was 2.6:1 to give the conditions described below and a residence time of 2,75 minutes was allowed in each mixer. The conditions were then as shown 55 in Table 6:—

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	TABLE 6				•		
5	System	Extraction pH	ra	lickel in affinate	NH ₃ in loaded organic phase gl ⁻¹		5
10	15% D2EHPA 4,4-4,5 15% D2EHPA 5% 2,8-3,2 0 EHO			,04 ,02	1,09 0,024		10
	A comparison of the D2EHPA-EHO system	acid-base ms given b	requiren elow in	nents for Table 7:	the D2EHPA and		
15	TABLE 7						15
			D2EHI	PA (%)	D2EHPA-EHO(%		
20	Ammonia for pH adjustment during extraction Acid make-up for return		125–133		85-98		20
	electrolyte Sodium hydroxide for adjustment of advance	r pH	25–3	33	16‡		
25	electrolyte Total acid-base		<1		5‡		25
	requirement		150-1	67	106-119		
30	the invention there	efore provid	des a sir	nple yet (s thus, to say the le effective process for exylic acids in comb	ast, appreciable. the extraction of metal ination with non-chelating	30
25	CLAIMAC						

35 CLAIMS

1. A process for the solvent extraction of metal values from solutions containing such metals comprising the contacting of a pregnant solution with a solvent extractant being an organophosphorus acid or a carboxylic acid capable of extracting desired metal values, the extraction being characterised by the fact that it takes place in the presence of a non-chelating oxime selected to 40 enhance extraction of the desired metal values.

2. A process as claimed in claim 1 in which the non-chelating oxime is selected to enable extraction to take place at a lower pH than was previously practiced.

3. A process as claimed in either of the preceeding claims in which the solvent extractant is an organophosphorus acid of the general formula (RO)₂PO-OH where each R is, independently 45 of the other, a substituted or unsubstituted alkyl, aryl or alkaryl group.

4. A process as claimed in any one of the preceeding claims in which the non-chelating oxime is one of an aldehyde where the alpha-carbon is primary or secondary.

5. A process as claimed in any one of the preceeding claims 1 to 3 and in which the nonchelating oxime is one of those specifically herein identified.

6. A process as claimed in any one of the preceeding claims in which the organophosphorus 50 or carboxylic acid is any one of those specifically identified.

7. A process as claimed in any one of the preceeding claims in which the process is one for the extraction of nickel, cobalt, manganese or copper. 8. A process as claimed in any one of the preceeding claims in which the extraction is

55 carried out at a pH of less than 5. 9. A process as claimed in claim 8 in which the extraction is carried out at a pH of less than

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10. A process substantially as herein described or exemplified.

11. Metals wherever separated by a method embodying the process of any one of claims 1 60 to 10. 60